

# **<sup>1</sup> Modeling the lava heat flux during severe effusive volcanic <sup>2</sup> eruption: an important impact on surface air quality**

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**Abstract.** The Reunion Island experienced its biggest eruption of Piton de la Fournaise volcano during April 2007. Known as “the eruption of the century”, this event degassed more than 230 KT of SO<sub>2</sub>. These emissions led to important health issues, accompanied by environmental and infrastructure degradations. This modeling study uses the mesoscale chemical model MesoNH-C to simulate the transport of gaseous SO<sub>2</sub> between April 2nd and 7th, with a focus on the influence of heat fluxes from lava. This study required the implementation of a reduced chemical scheme, a basic surface model and an estimation of lava heat fluxes in the atmospheric model. The model was able to reproduce general trends of this eruption, in particular the crossing of trade wind inversion, the SO<sub>2</sub> surface concentration (with highest peak of SO<sub>2</sub> of 600  $\mu\text{g m}^{-3}$  observed April 4th for western Reunion locations), and the wet deposition associated to rainfall. A sensitivity study shows that without heat fluxes over the vent and the lava flow, simulated SO<sub>2</sub> surface concentration are up to 45 times higher than observed.

## 1. Introduction

### 1.1. Generalities

Volcanoes are one of the most important natural sources of air pollution, both during and between eruptions [Oppenheimer, 2003]. It is essential for different areas of atmospheric science to have a good knowledge of volcanic volatile emissions in time and space, their atmospheric chemistry, physical and radiative effects. There are two different types of volcanoes: "reds" volcanoes characterized by relatively quiet effusive eruptions and transmitting any fluid lava in the form of castings, and "gray" volcanoes characterized by explosive eruptions and emitting pasty lava and ash in the form of pyroclastic flows. Each type of volcano is impacting the atmosphere in very different way, particularly in terms of injection depth and nature of the products ejected. Explosive volcanic eruptions such as those of El Chichon (Mexico) in 1982 [Pollack *et al.*, 1983; Hoffman, 1987] and Mount Pinatubo in 1991 [McCornick *et al.*, 1995; Fiocco *et al.*, 1996; Robock, 2002], mainly affected climate because of radiative and chemical impact of the plumes formed by aerosols injected into the stratosphere [Solomon, 1999; Robock, 2000, 2002]. For effusive volcanic eruptions such as those of Piton de la Fournaise (Reunion Island, Indian Ocean), the problem is different. Knowledge of their atmospheric and environmental impacts in the troposphere and degassing processes has some shortcomings. By chemical oxidation reactions, volcanic gases such as SO<sub>2</sub>, which is predominant during the degassing of the lava, become acidic and can interact with the aerosol phase as precursors of particles through nucleation and/or condensation. These tropospheric volcanic aerosols play an important role in atmospheric radiation, directly by scattering and absorbing of short wave radiation, and indirectly by changing cloud cover and cloud properties [Hobbs *et al.*, 1982; Al-

brecht, 1989; Kaufman et al., 2000; Yuan et al., 2011a, b]. Tropospheric volcanic aerosols and gaseous compounds, especially sulfur dioxide, can also be source of risks to terrestrial ecosystems and health at local or regional scales [Baxter et al., 1982; Mannino et al., 1996; Allen et al., 2000; Delmelle et al., 2001]. Piton de la Fournaise is a typical basaltic shield volcano located on the Indian Ocean Island of Réunion and, as Etna and Kilauea, it is one of the world's most active effusive volcanoes, with an eruption occurring every 10 months in average [Roult et al., 2012]. Reunion island is born 3 million years ago in the emergence of a gigantic volcano, in the southwest of the Indian Ocean at 21.06°S and 55.32°E. It presents on its 2512 km<sup>2</sup> a unique variety of landforms and landscapes with Piton des Neiges (3071m) being its highest point. The orographic influence on local dynamics of Reunion Island expected to be major. The interaction of the high mountainous terrain with the synoptic flow induces a large variability of wind field at local scale. The maritime and tropical location of the island, as well as the complexity of the terrain and wind exposure, imply a multitude of local circulations and weather, marked by large variations in temperature and precipitation. Modeling of local circulations is very complex to achieve in an environment like Réunion Island because it is the result of a complex interaction between topographic circulations, thermal breezes and local formation of clouds and precipitation. During an eruption, other parameters make modeling more difficult, including the effects of the dynamics of volcanic flows and thermal secondary effects associated with lava flow. Few studies, as the Vog Measurement and Prediction Project (VMAP) on island of Hawaii (<http://mkwc.ifa.hawaii.edu/vmap/hysplit>), have been able to accurately represent the distribution of volcanic pollution at characteristic scales of a volcanic island. The Piton de la Fournaise eruption of April 2007 presented all the characteristics of complex flow of sulfur dioxide with a temporal discontinuity between the highest concentrations of surface SO<sub>2</sub>

observed and the paroxysmal period of emission from the vent. The sulfur dioxide  $\text{SO}_2$  is the second gas emitted at the Piton de la Fournaise volcano after water  $\text{H}_2\text{O}$ , followed by carbon dioxide  $\text{CO}_2$  and hydrochloric acid  $\text{HCl}$ . The objectives in the framework of this case study are two-folds. First the paper aims to investigate the complex transport and distribution of the sulfur dioxide influenced by steep topography and three-dimensional atmospheric circulation. The second objective is to highlight the influence of the cloud scavenging and the heat fluxes on the  $\text{SO}_2$  surface concentration. For the latter, high resolution numerical simulations have been used to analyze the sensitivity of the heat fluxes on the volcanic pollutants. This paper starts with a brief description of April 2007 eruption as well as numerical methods (section 2). Section 3 is devoted to analyzing the estimation of the heat flux release during the eruption. The section emphasizes the influence of lava flow on the convection and its consequences on the  $\text{SO}_2$  distribution.

## 1.2. Description of the April 2007 eruption of the Piton de la Fournaise

In April 2007, the Reunion's Island has known its biggest eruption of Piton de la Fournaise volcano at least three centuries [Michon *et al.*, 2013]. Within a month,  $210 \text{ Mm}^3$  of lava flowed out with  $90 \text{ Mm}^3$  reaching the sea. Above all, the collapse of the summit caldera caused significant morphological change [Michon *et al.*, 2007]. Due to this nearby events and large environmental and civil protection impacts, this eruption is very well described in literature [Staudacher *et al.*, 2009; Vlastélic *et al.*, 2012; Barde-Cabusson *et al.*, 2011; Tulet and Villeneuve, 2010; Di Muro *et al.*, 2014]. After two short eruptive events (18 February and 30 March), a critical phase of the eruption started at 06 UTC on 2 April, located on the lower south-eastern part of the volcano ( $55^\circ 46' 25.5'' \text{ E}$  ;  $21^\circ 16' 54.6'' \text{ S}$  ,WGS84) at only 590m above the sea level and only 3km from the coast (Figure 1). In less than eleven hours, two main lava streams reached the sea,

83 producing significant water vapor plumes with a very low pH due to strong presence of sulfuric  
 84 and chlorohydric compound ( $\text{pH} < 2$ , *Staudacher et al.* [2009]). During the first two day, lava  
 85 fountains up between 50m and 150m high were observed. From 4 April, MODIS sensor shows  
 86 a significant increase of the lava flow rate until the 6th of April, where the peak of emissions of  
 87 lava was observed (greater than  $200 \text{ m}^3/\text{s}$  at the vent, *Coppola et al.* [2009]; *Staudacher et al.*  
 88 [2009]).  $\text{SO}_2$  emissions, proportional to lava emissions, were estimated at  $80 \text{ kg s}^{-1}$  the 4 April  
 89 12 UTC,  $320 \text{ kg s}^{-1}$  the 5 April 12 UTC and  $1600 \text{ kg s}^{-1}$  the 6 April 12 UTC before a strong  
 90 decrease until 8 April 12 UTC at  $55 \text{ kg s}^{-1}$ , and finally a constant emission of  $55\text{-}70 \text{ kg s}^{-1}$   
 91 until 11 April [*Tulet and Villeneuve*, 2010]. The peak of degassing was simulated at  $1800 \text{ kg}$   
 92  $\text{s}^{-1}$  on April 6, with the total budget estimated at 230kT, which is in agreement with the petro-  
 93 logic estimation of 311kT [*Di Muro et al.*, 2014]. The Observatoire Volcanologique du Piton de  
 94 la Fournaise (OVPF) recorded the 5 April at 20:48 UTC an earthquake of 4.8 magnitude syn-  
 95 chronous with the caldeira collapse [*Michon et al.*, 2007; *Staudacher et al.*, 2009]. As described  
 96 in *Tulet and Villeneuve* [2010] the location of this ash plume is well separate from the  $\text{SO}_2$  one,  
 97 as well as the vapor plume. The 6 April, lava fountains reached more than 200m high, and there  
 98 were several tens of individual lava flows from 2 to 20m wide [*Staudacher et al.*, 2009]. At this  
 99 moment, the lava flow reaches its maximum lateral and longitudinal extents. In late 6 April,  
 100 the intensity dramatically decreases, and the eruption became more “typical” compared to usual  
 101 eruptions of Piton de la Fournaise. The 12th of April, the shallow seismicity came back at its  
 102 highest, causing a new Dolomieu crater collapse. In the next days, the lava eruption intensity  
 103 became steady with effusion measured at  $15\text{-}20 \text{ m}^3.\text{s}^{-1}$ . This last event continued until May 1st  
 104 2007, the last eruption day.

### 1.3. ORA observations

ORA (Observatoire Réunionnais de l'air) provides daily monitoring of air pollution levels. It is equipped with several fixed and mobile stations along the Reunion coastline, measuring continuously primary and secondary pollutants. During the April 2007 eruption, 8 stations measured the SO<sub>2</sub> surface concentration from Saint Louis to Saint Denis passing through Cambaie in the west (Figure 1). Data from these three stations are compared to simulations. The first day, surface concentration of SO<sub>2</sub> is very low with value under 20  $\mu\text{g m}^{-3}$  for all stations (Figure 2). The next day from 02 UTC, Saint Louis and Cambaie stations measured significant increases of SO<sub>2</sub> concentrations with a peak of more than 60  $\mu\text{g m}^{-3}$  for Cambaie and 200  $\mu\text{g m}^{-3}$  for Saint Louis at 06 UTC. The 4th of April, new strong increases of surface concentration have been measured by all stations located on the southwest and northwest coast. A peak of 600  $\mu\text{g m}^{-3}$  was observed at Cambaie and 587  $\mu\text{g m}^{-3}$  in Saint Louis at 13 UTC. A significant decrease followed this peak of SO<sub>2</sub> surface concentration, with SO<sub>2</sub> concentration falling below the 100  $\mu\text{g m}^{-3}$  threshold at the end of the day. The 5th, a slight increase appears at dawn (from 03 UTC) with 200  $\mu\text{g m}^{-3}$  for Cambaie and 269  $\mu\text{g m}^{-3}$  for Saint Louis. Subsequently, from the 5th (12 UTC) to the 10th of April (06 UTC), the concentration varied between 20  $\mu\text{g m}^{-3}$  to 120  $\mu\text{g m}^{-3}$  for all west coast station except Saint Louis station, where brief peaks appeared for few hours on the 10th (345  $\mu\text{g m}^{-3}$ ) and the 24th (390  $\mu\text{g m}^{-3}$ ). The highest values measured by ORA are not in phases in time with the maximum emitted from the vent. This paradox needs a detailed study of sulfur dioxide transport.

## 2. Model description

### 2.1. Atmospheric model

The mesoscale non hydrostatic atmospheric model (MesoNH) developed by the Centre National de la Recherche Météorologique and the Laboratoire d'Aérodynamique [Lafore *et al.*, 1998] has been used for the study. MesoNH can be used at all scales ranging from synoptic to large eddy scales (<http://mesonh.aero.obs-mip.fr/>). It can be run in a two way nested mode involving up to eight nesting stages. Different sets of parameterizations have been introduced for convection [Bechtold *et al.*, 2001; Pergaud *et al.*, 2009], cloud microphysics [Cohard and Pinty, 2000], turbulence [Bougeault and Lacarrere, 1989], lightning [Barthe *et al.*, 2007], gaseous chemistry [Suhre *et al.*, 1998; Tulet *et al.*, 2003], cloud chemistry [Leriche *et al.*, 2000] and aerosols [Tulet *et al.*, 2005; Grini *et al.*, 2006].

### 2.2. Surface model

The SURFEx surface scheme is coupled with MesoNH to simulate surface processes, thermodynamic and chemical exchanges with the atmosphere. (<http://www.cnrm.meteo.fr/surfex/>; Masson *et al.*, 2013.). SURFEx is composed by various parameterizations for natural land surface [Noilhan and Mahfouf, 1996; Bougeault and Lacarrere, 1989], urbanized area [Masson, 2000], lakes and oceans [Salgado and Le Moigne, 2010] and chemistry and aerosols surface processes [Tulet *et al.*, 2003; Mokhtari *et al.*, 2012]. The coupling with the atmospheric model is performed by averaging the surface fluxes over a model grid box. Within SURFEx, the lava flow is represented by a line of potential heat flux emission, starting from the vent of the volcano at 21.28°S and 55.77°E to the coastline at 21.28°S and 55.80°E. This representation implies two major approximations for the lava flow. The first one is the misrepresentation of lava flow shape, as the observed lava flow has a triangular shape. The second is relative to the static represen-



tation of the lava under SURFEx, when the lava propagation is not integrated in time. The increasing surface of lava flow and its heat flux is modeled by multiplying the line of potential emission with a coefficient proportional to the increase of the lava flow surface.

### 2.3. Model configuration

The simulation starts at 00 UTC on April 2nd 2007, and ends at 00 UTC on April 7th 2007. The simulation has two nested domains with Kessler microphysics scheme and TKE turbulence scheme (prognostic turbulent kinetic energy, one and a half order closure). The largest domain with high model grid spacing (2km) is centered over the Reunion island. The domain extends over 330km from north to south and 450km from east to west. The second domain covers only the Reunion Island and its coastline with a horizontal model grid spacing of 500m. The vertical grid is composed of 72 levels for both models stretching up to 31km altitude with a first level 5m above ground level. Initial and lateral boundary conditions are extracted from ECMWF analysis for the meteorological fields and from MOCAGE (<http://www.cnrm.meteo.fr/gmgec/spip.php?article87>) for gaseous chemistry fields. The gas phase chemistry is resolved on both domains using the ReLACS chemical mechanism [Crassier *et al.*, 2009], which is a reduced version of RACM including the oxidation of sulfur dioxide by OH radical. In SURFEX, the entire SO<sub>2</sub> emission is released at the vent. This assumption is well correlated with the fact that the magma begin to degas when it reaches the vent and its surroundings. In consequence, with a 500m MesoNH horizontal model grid spacing, the location of SO<sub>2</sub> emission is well represented. A simulation protocol was implemented to limit the model drift by reinitializing the model dynamic (wind, humidity and temperature filed) in the middle of the simulation while the chemical fields have been preserved along the whole period (Figure 3). A second simulation starts the 3rd at 18 UTC until the 4th of April at 00 UTC. This latter

provides at its end the dynamic fields refresh, while suppressing the need for model spin-up (time taken by the model to reach equilibrium state). Finally, a new simulation starts the 4 April at 00 UTC with the dynamic of simulation 2 and chemistry of simulation 1.

Sensitivity tests are made in this study to highlight the influence of heat fluxes in the transport of volcanic pollutants and the influence of cloud chemistry in scavenging sulfur dioxide. The 3 simulations configurations are sum up in the table 1.

### 3. Estimations of thermodynamic emissions

#### 3.1. Heat flow estimation

Lava heat is released in the atmosphere from the core of an active flow by conduction through the basal, lateral and surface crusts [Oppenheimer, 1991; Klingelhofer *et al.*, 1999; Quarenii *et al.*, 2004]. At the surface, heat losses are dominated by radiation ( $5 \times 10^4 \text{ W m}^{-2}$ ) and convection ( $10^4 \text{ W m}^{-2}$ ), whereas conduction from the base to the ground is predominant ( $10^3 \text{ W m}^{-2}$ , Harris *et al.* [2005]). For this study, only convective heat fluxes are implemented, with the assumption that heat losses by conduction and by rain falling on the flow ( $250 \text{ W m}^{-2}$ , Harris *et al.* [2005]) are negligible. As the influence of radiant heat fluxes is inversely proportional to the square of the distance; we have also neglected it for our simulation. The heat flow by convection is calculated from:

$$Q_{conv} = hc(T_{surf} - T_{air})$$

With  $hc$  the heat transfert coefficient estimated at  $50 \text{ W m}^{-2}$  by Keszthelyi *et al.* [2003],  $T_{surf}$  the lava surface temperature and  $T_{air}$  the air temperature (290K). Estimation of the sensible heat fluxes, and hence the lava cooling, is mainly controlled by the surface winds. The heat flux relation to the wind from Keszthelyi observations are taken into account in our model (Figure

185 4). The surface covered by hot (1100°C) liquid lava flow and the crusted (400°C) lava is taken  
 186 from the day by day observation given by *Bachèlery et al.* [2014] between April 2nd and 8th,  
 187 allowing the estimation of the heat flow (in  $m^2$ ) from the ground.

#### 4. Sulfur transport during the April 2007 eruption of Piton de la Fournaise volcano using MesoNH atmospheric model

##### 4.1. SO<sub>2</sub> mass burden

188 Figure 5 represents the evolution of SO<sub>2</sub> mass burden simulated by MesoNH between April  
 189 3rd and April 6th at 18 UTC. The first period until 4 April shows that the plume is oriented to  
 190 the west with a maximum of mass burden of 210 DU. From April 5 and 6, the plume at the  
 191 vent, above 5km ASL is oriented to the north with a large value (330 DU) over Reunion island.  
 192 During this period, the strong presence of SO<sub>2</sub> in the north of Reunion Island indicates a plume  
 193 separation. The change in direction between the two periods is due to the SO<sub>2</sub> plume crossing  
 194 the inversion of trade winds around 3500m-4500m ASL. Below the inversion, a lower branch  
 195 of the SO<sub>2</sub> plume is transported westward by the trade winds, while above, an upper plume is  
 196 advected eastward. This analysis of SO<sub>2</sub> plume evolution in the first 4 days of the eruption is  
 197 consistent with the study based on satellite data OMI and CALIOP by Tulet and Villeneuve  
 198 (2010).

##### 4.2. Simulated SO<sub>2</sub> surface concentrations

199 In the morning of April 2, the SO<sub>2</sub> plume at the surface is oriented southwestwards, contourn-  
 200 ing by the south the Piton de la Fournaise area. The plume is then transported along the coastline  
 201 influenced by the trade winds circumventing the island, where a strong gradient of SO<sub>2</sub> appears.  
 202 During this day, the SO<sub>2</sub> plume reaches Cambaie, at the northwest of the island with low surface  
 203 concentration of SO<sub>2</sub> in order of some tens of  $\mu g m^{-3}$ . The strongest simulated concentrations

are located at low altitude, near Saint Joseph in the south, with  $350 \mu\text{g m}^{-3}$ . For other stations,  $\text{SO}_2$  surface concentration is  $125 \mu\text{g m}^{-3}$  at Saint Louis in the southwest or a comparatively low  $25 \mu\text{g m}^{-3}$  is observed in Sainte Thérèse in the northwest, while concentrations in Saint Denis to the north is even lower with  $1 \mu\text{g m}^{-3}$ . Not much changes occur the 3th of April, when the overall atmospheric dynamics confine the volcanic pollutants to the west of the island (Figure 6, dots represent ORA stations, Saint Denis at the north, Cambaie at the northwest and Saint-Louis at the southwest). Concentrations for the majority of the stations are stronger than the eve, with  $200 \mu\text{g m}^{-3}$  at Saint Louis and  $75 \mu\text{g m}^{-3}$  at Cambaie and Sainte Thérèse (Figure 6). However, the strong  $\text{SO}_2$  gradient noted earlier is no longer present on April 3rd.  $\text{SO}_2$  distribution appears larger on the south and west side of La Reunion and to the west above the ocean.

Higher concentrations are simulated the 4th of April, with a peak of  $680 \mu\text{g m}^{-3}$  obtained at St. Joseph and  $350 \mu\text{g m}^{-3}$  obtained at St. Louis. For stations in the north of the island, the concentrations are close to of those April 3rd. The maximum  $\text{SO}_2$  concentration at the surface is thus located in the heights of the island, where high concentrations are simulated notably with more  $1000 \mu\text{g m}^{-3}$  in the heights above the city of Saint Joseph (south of Réunion Island). On the 5th of April, an increase of heat flow and a decrease in the stability of the atmospheric boundary layer allows the plume to reach higher altitude and in consequence be oriented directly to the northwest.

Finally, the 6 April, high concentrations are only simulated over the entire southern half of the island, with concentrations over  $5000 \mu\text{g m}^{-3}$ , 10 times the threshold recommended by European standards. Unfortunately, no observations are available in the south of the island to validate these very high-simulated concentrations the 6 April.

### 4.3. Comparison between MesoNH simulation and ORA data

From 2 to 5 April, the simulation succeeds to correctly reproduce general trends for all simulated stations on the island (Figure 7). On the 2nd, the  $\text{SO}_2$  surface concentration given by MesoNH corresponds to ORA observations for these three stations with values below  $15 \mu\text{g m}^{-3}$  for Cambaie and Saint Denis and a peak of  $200 \mu\text{g m}^{-3}$  in the middle of the day for Saint Louis. On the 3rd, Saint Denis station did not record any presence of  $\text{SO}_2$  while Cambaie station had a gradual increase with a peak of  $85 \mu\text{g m}^{-3}$  for ORA observation and  $135 \mu\text{g m}^{-3}$  in MesoNH simulation. Saint Louis has experienced a significant increase with a  $\text{SO}_2$  surface concentration of  $480 \mu\text{g m}^{-3}$  measured by ORA and of  $605 \mu\text{g m}^{-3}$  simulated by MesoNH, once again in the middle of the day. The 4th of April, no changes occurred for Saint Denis, while a strong  $\text{SO}_2$  increase appears at Cambaie with  $500 \mu\text{g m}^{-3}$  in MesoNH and  $601 \mu\text{g m}^{-3}$  for ORA observations. The same behaviour is also seen for Saint Louis station, with  $585 \mu\text{g m}^{-3}$  observed against a strong over concentration simulated value of  $1135 \mu\text{g m}^{-3}$ . It is important to note that this latter station is positioned on a very strong gradient of  $\text{SO}_2$  ( $1135 \mu\text{g m}^{-3}$  to  $220 \mu\text{g m}^{-3}$  at 5km away). This strong increase is immediately followed by a sharp decrease at the end of the day, with value below  $150 \mu\text{g m}^{-3}$  for Cambaie and  $200 \mu\text{g m}^{-3}$  for Saint Louis. On the 5th  $\text{SO}_2$ , concentration varies between 30 to  $200 \mu\text{g m}^{-3}$  for Cambaie, and 10 to  $300 \mu\text{g m}^{-3}$  for Saint Louis except in the evening (18 UTC) where the simulation does not succeed to keep low concentration values ( $550 \mu\text{g m}^{-3}$  simulated instead of  $35 \mu\text{g m}^{-3}$  observed). For the simulation's last day, the  $\text{SO}_2$  surface concentration given by the model is stronger than observations, with highest values of more than  $450 \mu\text{g m}^{-3}$  (against  $45 \mu\text{g m}^{-3}$  observed) at Cambaie and a peak of  $590 \mu\text{g m}^{-3}$  at Saint Louis instead of  $55 \mu\text{g m}^{-3}$ . The same anomaly appears for Saint Denis, when shortly after 00 UTC, the simulation gives a peak of  $450 \mu\text{g m}^{-3}$  instead of a total absence of

248 volcanic SO<sub>2</sub> highlighted by ORA measurements. Despite these orders of magnitude anomalies  
 249 from the April 6th, and the global over exposition of SO<sub>2</sub> surface concentration for Saint Louis,  
 250 SO<sub>2</sub> concentrations between 2 and April 7 are generally consistent with ORA measurements.  
 251 The simulation succeed to recreate the paradoxical situation between the highest surface SO<sub>2</sub>  
 252 concentration measured by ORA the 4 April whereas the paroxysmal intensity of the eruption  
 253 is in the night (from 20 UTC) of the 5 April and on April 6th.

#### 4.4. Vertical transport above the eruption

254 The increased lava flow and its greater surface coverage between April 2 and 7 consequently  
 255 involves an increase in heat flux over the lava flow. Heat flow for the first three days are moderate  
 256 with an average of 12800 W m<sup>-2</sup>. Local circulation is still dominated by trade winds with  
 257 surface winds around 5 m s<sup>-1</sup>. The plume reached 2.5km ASL. (under 100 μg m<sup>-3</sup>) and the  
 258 highest SO<sub>2</sub> concentration value are close to the surface (23000 μg m<sup>-3</sup>). However, from April  
 259 5th, the general trend is the increasing of trade winds (11 m s<sup>-1</sup> around the eruption zone). This  
 260 increase induces more heat flux (22500 W m<sup>-2</sup> April 5) and a local breeze in the eruption area  
 261 which creates a more efficient vertical transport of volcanic sulfur.

262 The surface warming and the heat flow associated with the lava flow generate atmospheric  
 263 instability in the low layers of the troposphere. On Figure 8, the strong convection above the lava  
 264 flow creates a large mixing area with a maximal negative vertical gradient of equivalent potential  
 265 temperature of  $\partial\theta_e/\partial z = -1.5\text{K/km}$  between the lava and 7km ASL. Under the influence of the  
 266 trade winds, the vertical structure of the plume in altitude is moving slightly westward. The  
 267 vertical wind above the eruption reached 14 m s<sup>-1</sup> from 3 to 5km ASL. and transports SO<sub>2</sub> up  
 268 to 8 km high (under 100 μg m<sup>-3</sup>), ie above the inversion zone trade winds situated between 2.5  
 269 and 3.5 km of altitude. At this altitude the plume is in thermodynamic equilibrium with the

environment and oriented according to the wind direction, ie west/southwest. The plume is no longer transported to the west, but in the direction of Mauritius and Australia. High values of  $\text{SO}_2$  concentration are modeled up to 6km (above  $30000 \mu\text{g m}^{-3}$ ) the 6 April (Figure 9, cross section in left panel corresponds to blue line (2 April) in Figure 1 and cross section in right panel corresponds to red line (6 April) ).

#### 4.5. Rainfall and aqueous chemistry

The 5 and the 6 April, strong clouds formations appears in the Piton de la Fournaise area. The associated accumulated rainfall simulated between 2 and 7 April 2007 by MesoNH is consistent with Meteo-France observations (Figure 10). Only the southeast weather station gives high rainfall value (67mm cumulated), whereas the western weather station recorded lower value (15mm and 6mm). One possible causes of concentrations overprediction for April 6th is that the scavenging of  $\text{SO}_2$  by rain and cloud water leading to sulfuric acid formation is not taken into account in the simulation. However, Meteo-France measurements have shown that between April 2 and 7, the largest quantity of rain were observed only for the 6 April in the volcanic region. As MesoNH includes a cloud chemistry module [Leriche *et al.*, 2013] a sensitivity test was realized from April 5 18 UTC to April 7 00 UTC (limited period due to high computational cost) by activating this module. A simplified mechanism in aqueous phase was used including the oxidation of  $\text{SO}_2$  into sulfuric acid by hydrogen peroxide, ozone and pernitric acid [Leriche *et al.*, 2003]. The module includes also the mass transfer kinetic for the exchange between the gas phase and liquid phases of soluble gases and their redistribution between cloud water and rainwater by microphysical processes (collision/coalescence leading to precipitation and sedimentation of raindrops leading to wet deposition).

One of the main consequence of cloud chemistry activation is a global decrease of  $\text{SO}_2$  surface concentration due to  $\text{SO}_2$  scavenging by rainfall and aqueous phase  $\text{SO}_2$  chemistry transformation inside clouds over Reunion Island. The 6 April at 13 UTC, the difference of  $\text{SO}_2$  surface concentration between the simulation with cloud chemistry activated (AQ simulation) and reference simulation (REF) reach up to  $-700 \mu\text{g m}^{-3}$  over the high terrain in the center of the island (Figure 11). The  $\text{SO}_2$  surface concentration for the western coastline is  $200 \mu\text{g m}^{-3}$  lower for the AQ simulation than the REF simulation and for the Piton de la Fournaise area, a strong decrease appears due to the proximity of the vent with the presence of high rainfall this day over the volcano. Generally, a 30% to 60% decrease is simulated by MesoNH, giving  $\text{SO}_2$  surface concentrations close to ORA observations (Figure 12).

## 5. Influence of sensible heat fluxes in the transport of $\text{SO}_2$

A sensitivity study was performed to characterize the influence of heat flux forcings over the vent and lava on the vertical transport of  $\text{SO}_2$  (Figure 9). To do so, an additional simulation has been made without thermodynamic flux (NO-FLX) to highlight the contribution of these fluxes in the transport and the dispersion of sulfur dioxide. This simulation rapidly presents large discrepancy from the reference simulation (REF) as shown by Figure 13 in the differences of  $\text{SO}_2$  concentration (in  $\mu\text{g m}^{-3}$ ) between the NO-FLX simulation and the REF simulation on the 3,4,5 and 6 of April at 13 UTC. A strong positive difference in concentrations appears for the whole southern part of the island with a maximum of  $32000 \mu\text{g m}^{-3}$ . The northwestern part of the island is also overexposed to higher concentrations of the order of  $500 \mu\text{g m}^{-3}$  for April 3 and  $1000 \mu\text{g m}^{-3}$  from April 3 to 5. Conversely, negative anomalies are simulated the 4th of April for the northwest with less than  $340 \mu\text{g m}^{-3}$  compared to concentrations in the REF simulation. As a main consequence, concentrations obtained with the simulation NO-FLX are



also far from ORA measurements. In general, unrealistic peaks are simulated (Figure 14) with a factor of 5 to 35 in the south, and 5 to 10 in the northwest compared to ORA observations. Taking into account sensible heat flux from lava is therefore of prime importance as the NO-FLX simulation did not recreate correctly the spatial and temporal distribution of sulfur dioxide for the 2007 eruption of Piton de la Fournaise. Here, the lack of heat flux injection did not allow adequate vertical transport, essential to get an overall good representation of SO<sub>2</sub> distribution. As a general consequence we estimate that numerical modeling of the April 2007 eruption cannot be represented without heat flux correctly estimated and injected at the eruptive vent.

## 6. Conclusions

The objective of this study was to model fine scale spatial distribution of SO<sub>2</sub> degassed during the eruption of the Piton de la Fournaise in April 2007. It was necessary to adequately modeled the heat flux injection over the vent and lava flow to simulate the atmosphere dynamics that drives this SO<sub>2</sub> distribution. The simulation has been found to be in relatively good agreement with observations, and highlighted two phases. With moderate value of heat fluxes from lava flow (12800 W.m<sup>-2</sup>), the first phase, between April 2 and 4, shows a SO<sub>2</sub> plume still contained under the trade wind inversion at 3km ASL. The main consequence is a high SO<sub>2</sub> surface concentration for western stations (600 μg m<sup>-3</sup> for ORA observations, 500 μg m<sup>-3</sup> for simulation at Cambaie). The second phase, between April 5 and 7, corresponds to the eruption maximum intensity. This high intensity is accompanied by a strong increase in lava heat flux (22500 W m<sup>-2</sup>) that allows the SO<sub>2</sub> plume to cross the trade wind inversion, and reach an altitude of 8km ASL on the 6th of April. This deep convection reduces surface SO<sub>2</sub> concentration (600 μg m<sup>-3</sup> to 100 μg m<sup>-3</sup> in few hours at Cambaie), but the model fails to keep low SO<sub>2</sub> surface concentration on the last simulation day (peaks at 400 μg m<sup>-3</sup> instead of 55 μg m<sup>-3</sup> at the end of 6 April). These

over predictions were addressed by taking into account the cloud chemistry in a sensitivity study realized from 12 UTC on April 5 to April 7. During this period, the scavenging of  $\text{SO}_2$  by rain water and cloud water significantly reduces  $\text{SO}_2$  surface concentration, producing sulfuric acid and as a consequence acid rain. Overall, the reference simulation was largely in good agreement and within the same order of magnitude, with the observation values from ORA. To highlight heat flux influence, a second sensitivity study was performed, in which the heat fluxes from the vent and lava flow were totally suppressed. Without these additional contributions of heat flux, the simulated surface concentrations are up to 45 times higher than the observations. One of the main conclusions of the study is that heat flux above lava is a crucial parameter to take into account in order to reproduce correctly  $\text{SO}_2$  distribution. This additional energy allows the development of strong convection that injects volcanic discharges over the atmospheric boundary layer. The heat flux model, although still imperfect by its surface representation, significantly improve the  $\text{SO}_2$  spatial distribution, as shown in this study by respecting orders of magnitude compared to observations and by displaying correct temporal evolution of the simulated surface concentrations.

A perspective of improvement is the implementation of a new deep convection scheme to improve the representation of sub-grid convective transport in MesoNH model. The initial deep convection scheme from MESO-NH basic package is not adapted for an extreme event such as volcanic eruption. Indeed, some important processes are not taken into account or are not representative of a phenomenology of an eruption, such as the speed of ejection of gas and heat flow, or the absence of the mixing vertical processes. A strategy could be a coupling system between a more detailed lava surface model and MesoNH atmospheric model to better reproduce the distribution and evolution of the lava during the period.

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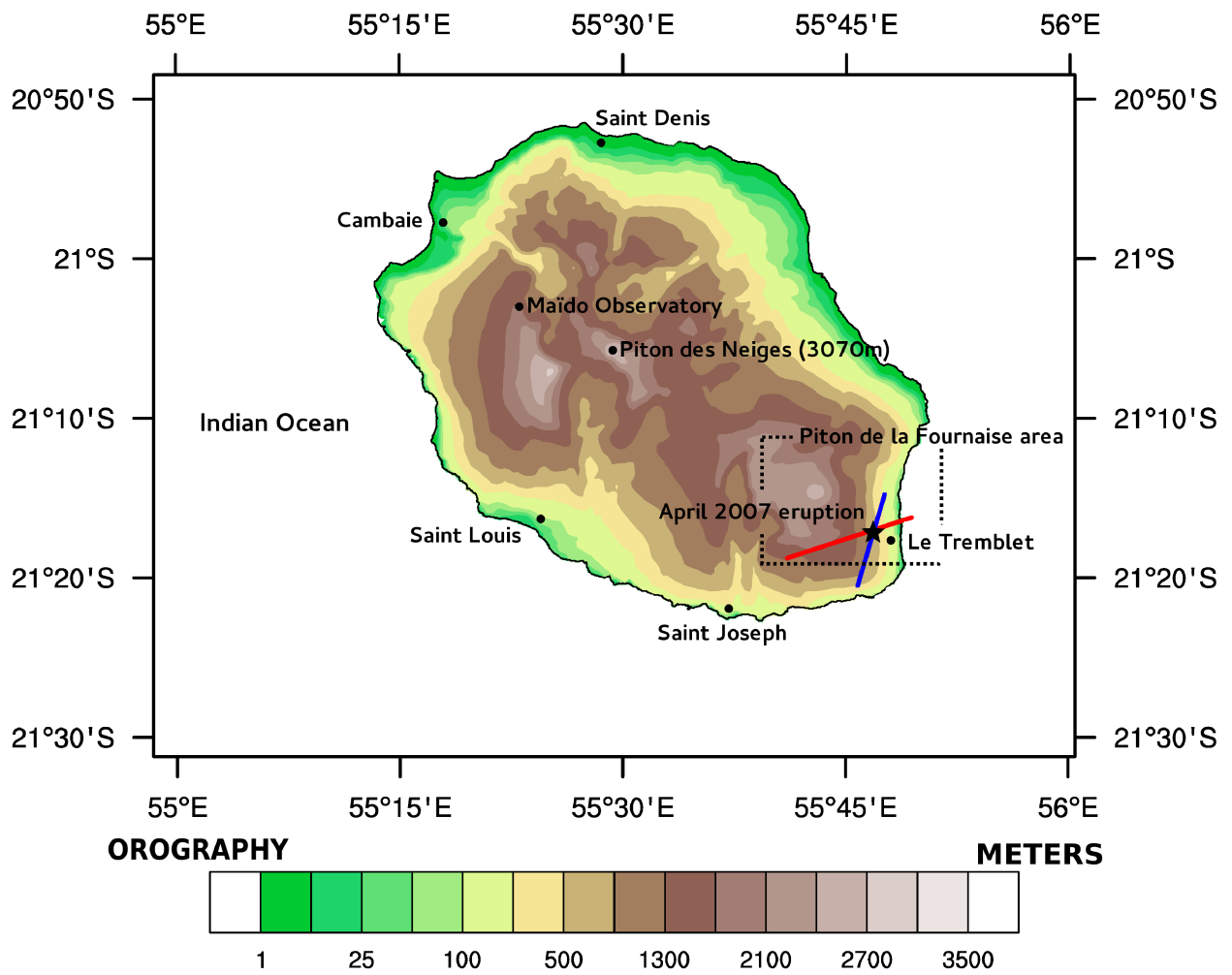


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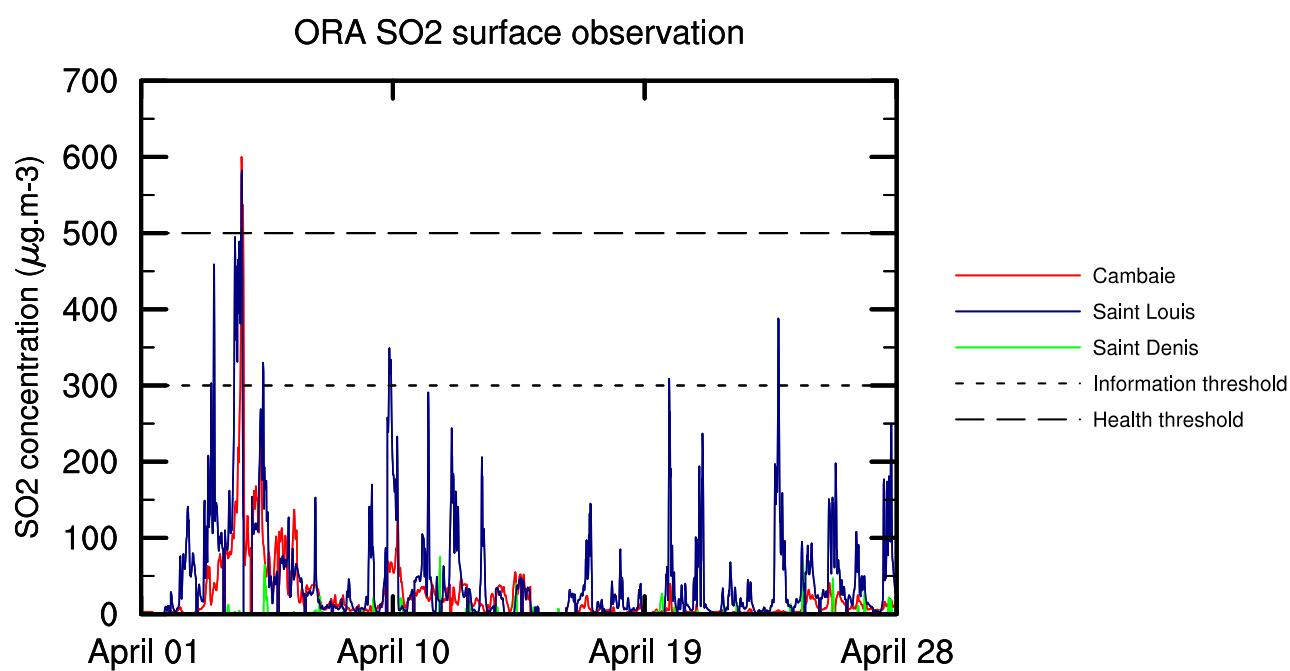
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Simulation	Period	Lava heat flux	Cloud chemistry
REF	04/02 00 UTC - 04/07 00 UTC	Yes	No
NO-FLX	04/02 00 UTC - 04/07 00 UTC	No	No
AQ	04/05 18 UTC - 04/07 00 UTC	Yes	Yes

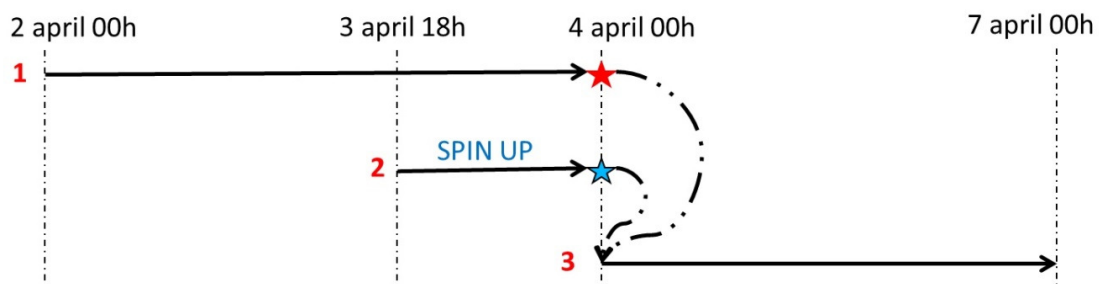
**Table 1.** The 3 simulations configurations. REF is the reference simulation, NO-FLX is the simulation without heat fluxes from lava flows and AQ is the simulation with cloud chemistry activated



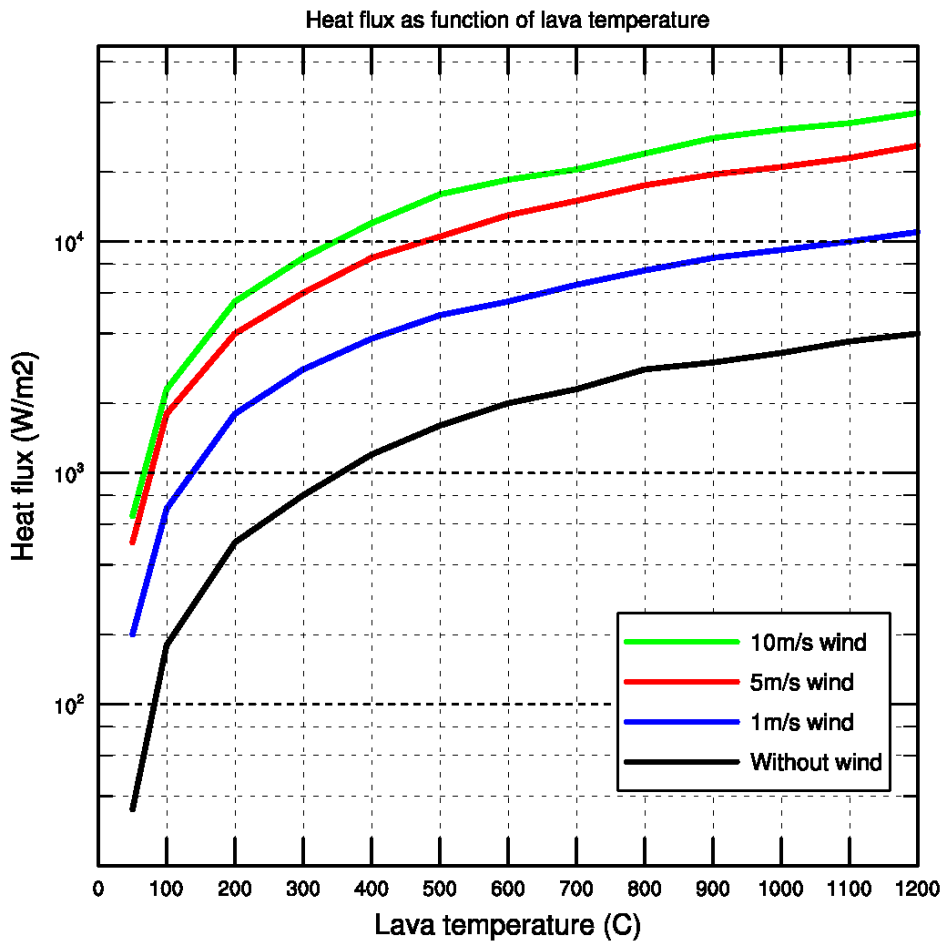
**Figure 1.** Orography and geographic situation of Reunion Island. The blue line and the red line in the Piton de la Fournaise area correspond respectively to the cross section of 2 April 2007 and 6 April 2007 describes in section 4-3.



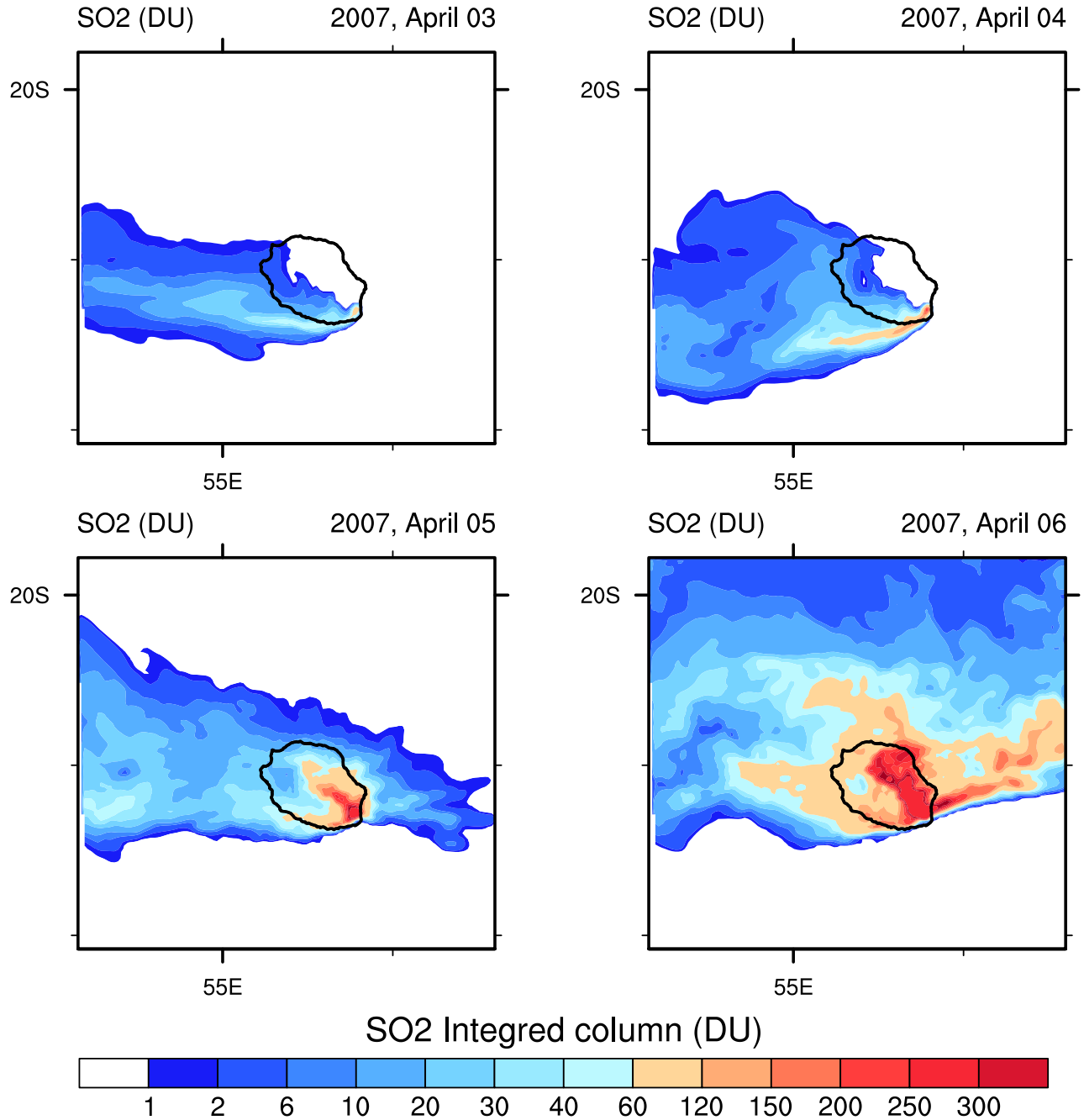
**Figure 2.** ORA measurements between April 1st and April 28th for Cambaie in the northwest (red), Saint Louis in the southwest (blue) and Saint Denis in the north (Green). Thin dashed line is the public information threshold and the large dashed line is the health threshold.



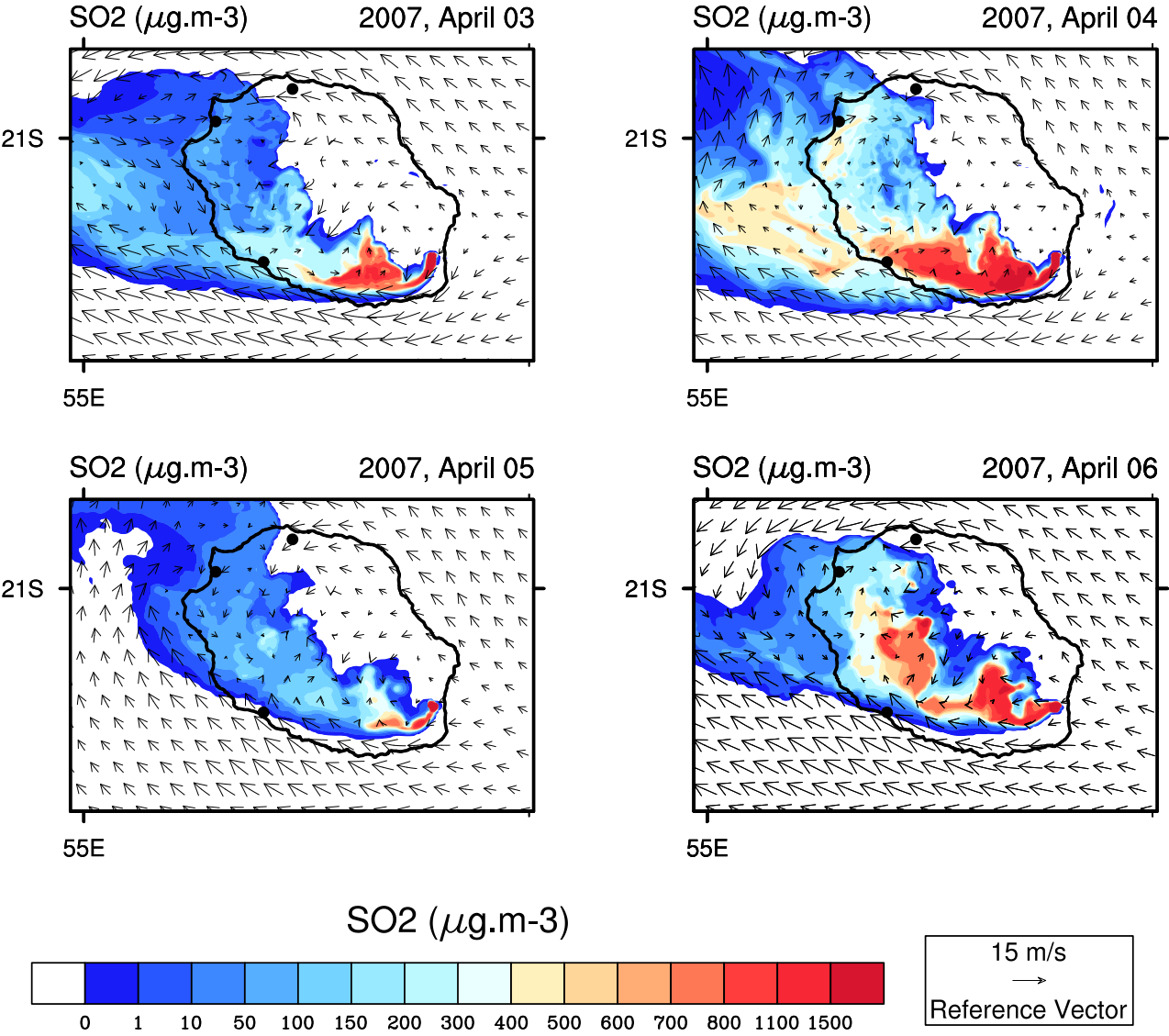
**Figure 3.** Updating the model dynamic: As a first step, the reference simulation begins the 2 April until 4 April 00 UTC (1). Then a new simulation begins April 3 18 UTC until 4 April 00 UTC (2). This latter will give the new model dynamics while avoiding the early simulation spin up. Finally, the REF simulation resumes the 4 April 00 UTC into the end, with chemical fields of (1) and model dynamic of (2).



**Figure 4.** Heat flux evolution with lava surface temperature (Kezsthelyi et al, 2003). The green, red, blue and black lines represent heat fluxes respectively for  $10 \text{ m s}^{-1}$ ,  $5 \text{ m s}^{-1}$ ,  $1 \text{ m s}^{-1}$  and without surface wind.

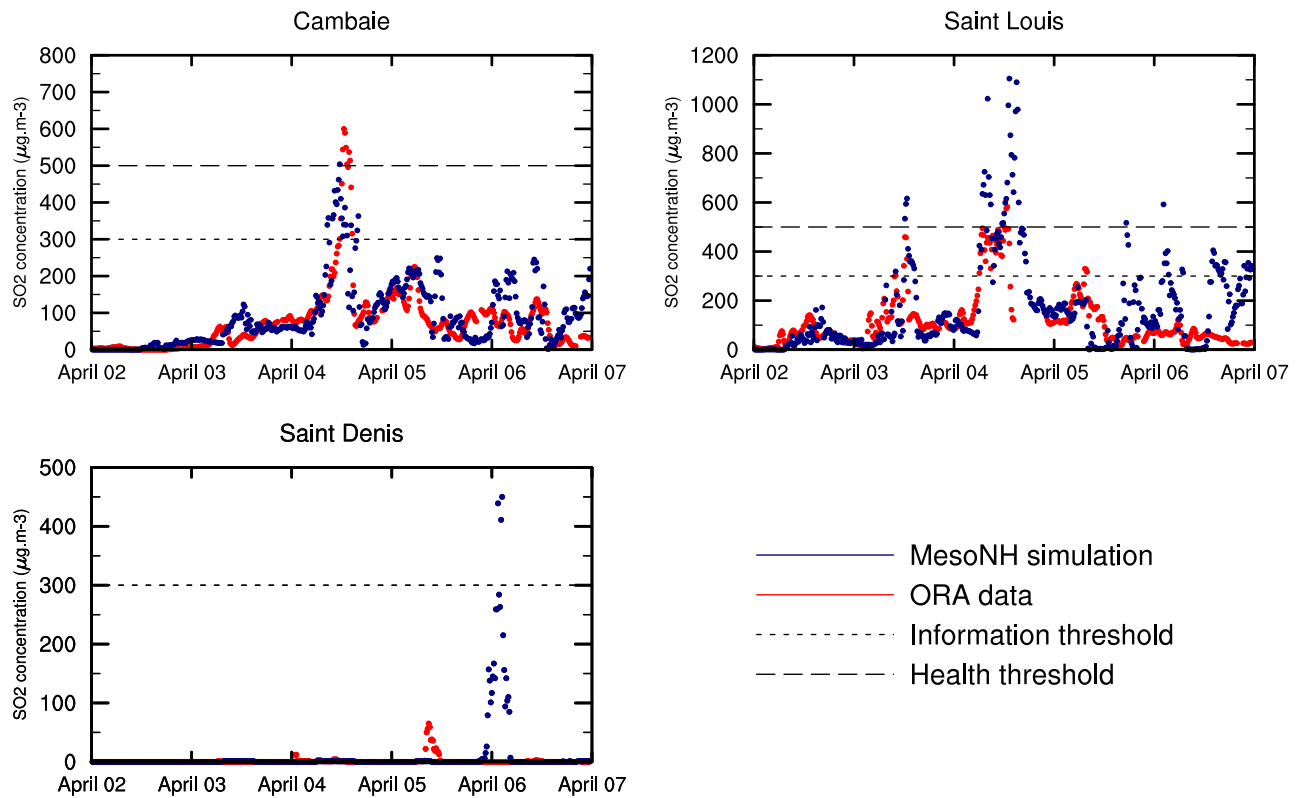


**Figure 5.** Integrated column of SO<sub>2</sub> (DU) between April 3 and April 6 at 13 UTC above the Reunion Island from first model domain (2km horizontal model grid spacing ). April 2 and 3 , the SO<sub>2</sub> plume, influenced by the trade winds below the thermic inversion, is oriented to the west. The 4 and 5 April, a large part of the SO<sub>2</sub> plume are crossing the trade winds inversion and is transported to the northeast.

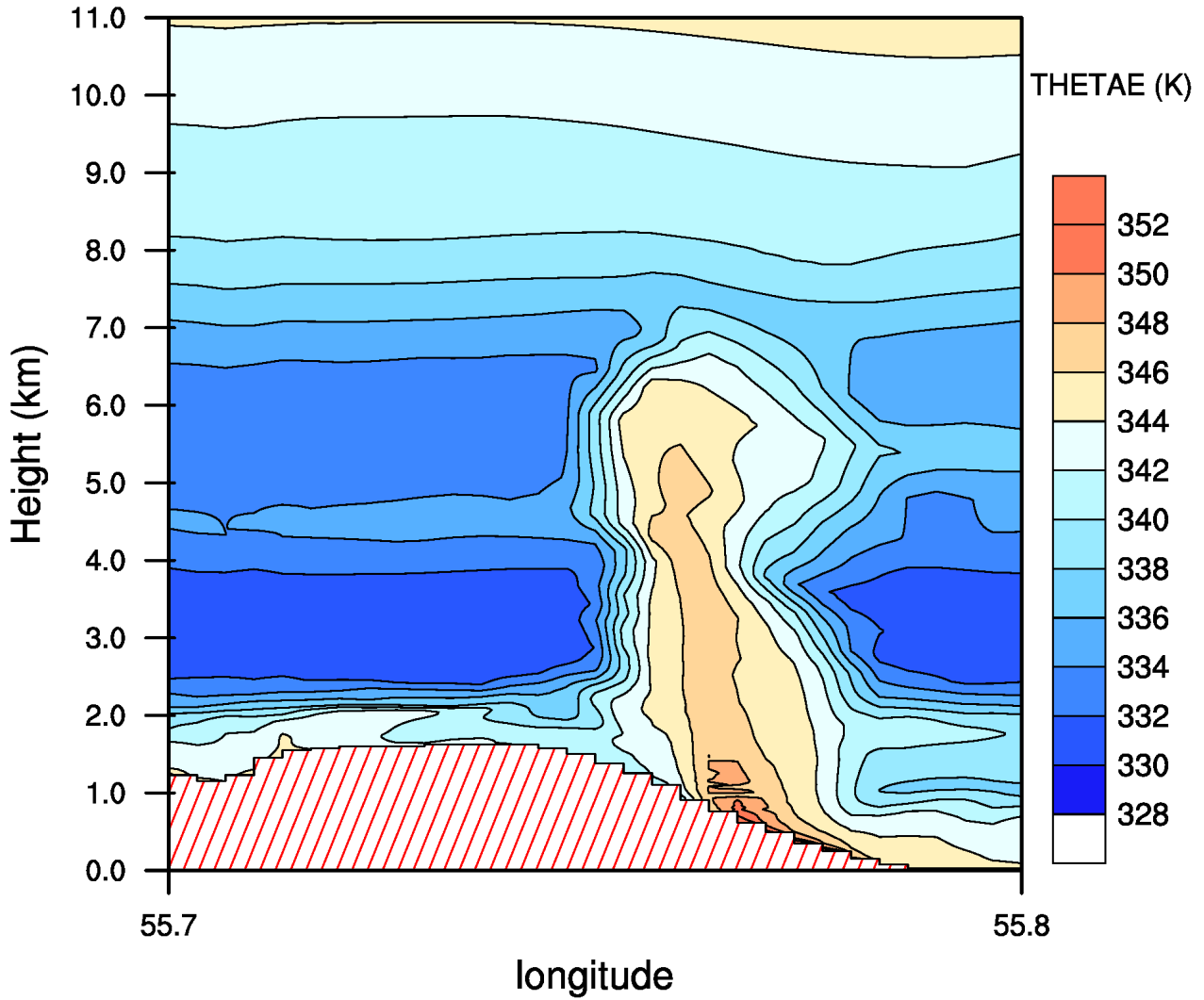


**Figure 6.** Surface concentration of SO<sub>2</sub> between April 3 and 6 at 13 UTC from MesoNH mesoscale atmospheric model smallest domain (500m horizontal model grid spacing)

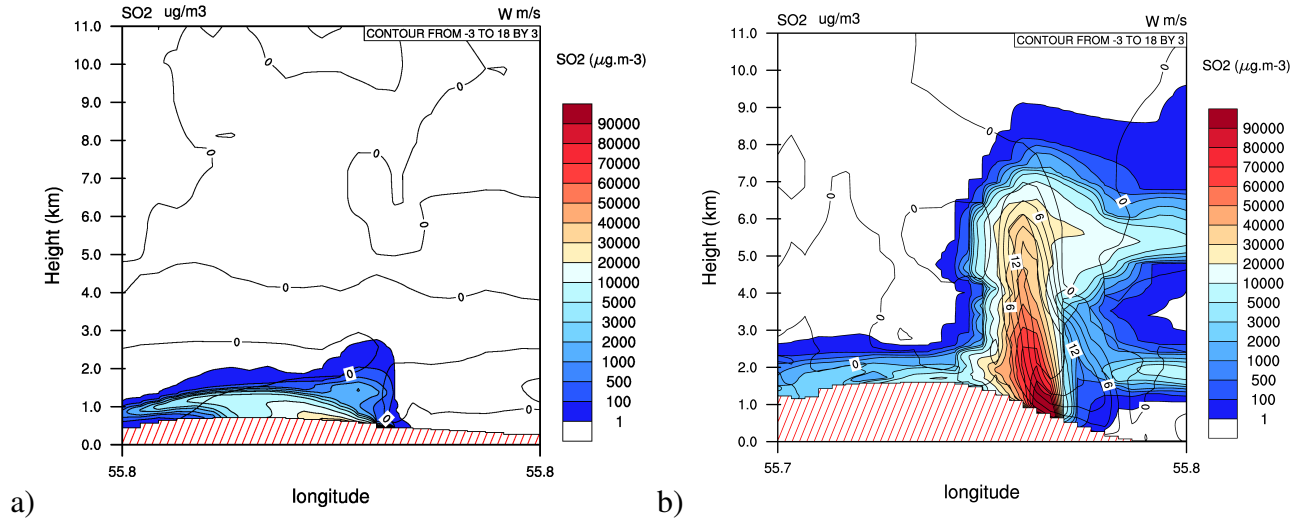




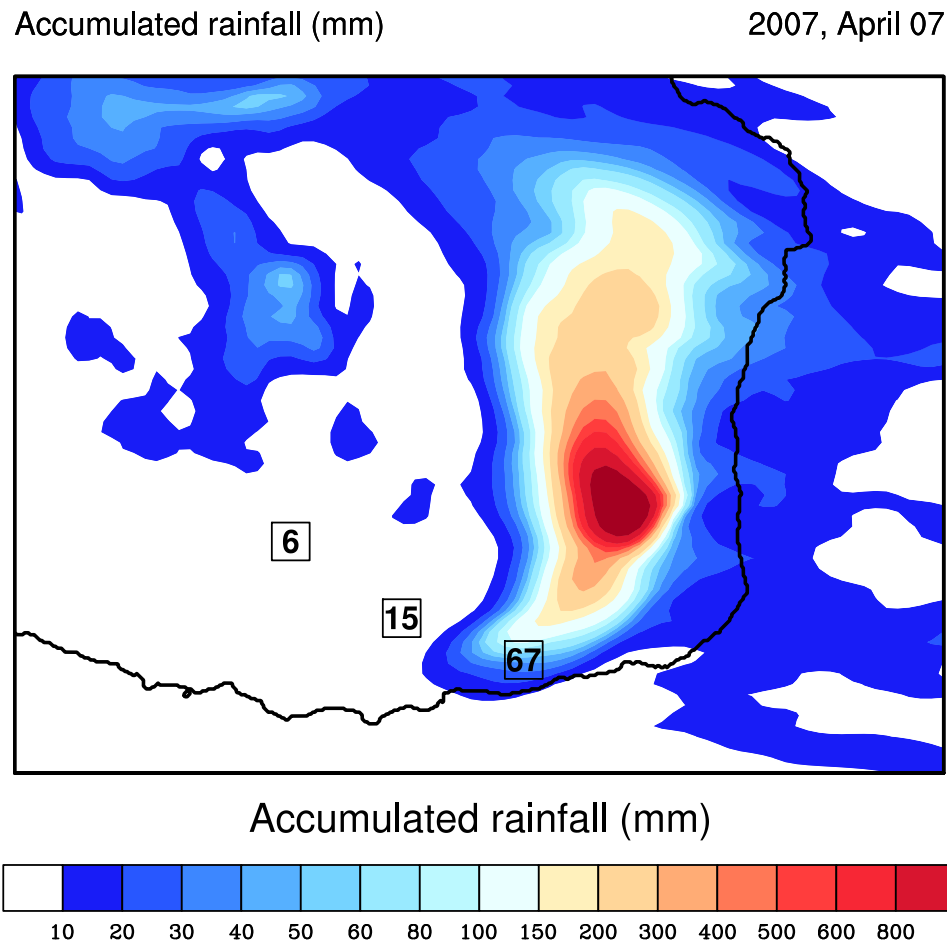
**Figure 7.** Comparison between ORA observation (blue points) and MesoNH simulation (red points) from April 2 to 7 April. The large dashed line is the health threshold while the thin dashed line is the information threshold.



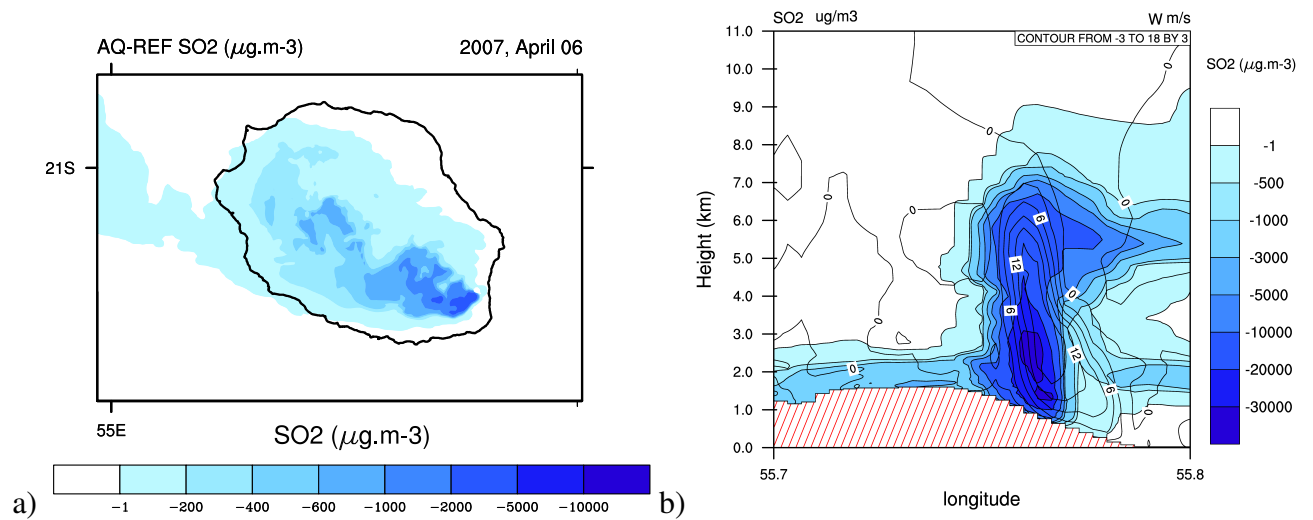
**Figure 8.** Cross section of equivalent potential temperature (K) for the 6th April at 13 UTC along red line in Figure 1. The strong convection above the lava flow creates a large mixing area with a maximal negative vertical gradient of equivalent potential temperature of  $\partial\theta_e/\partial z = -1.5\text{K/km}$  between the lava and 7km ASL. Under the influence of the trade winds, the vertical structure of the plume in altitude is moving slightly westward.



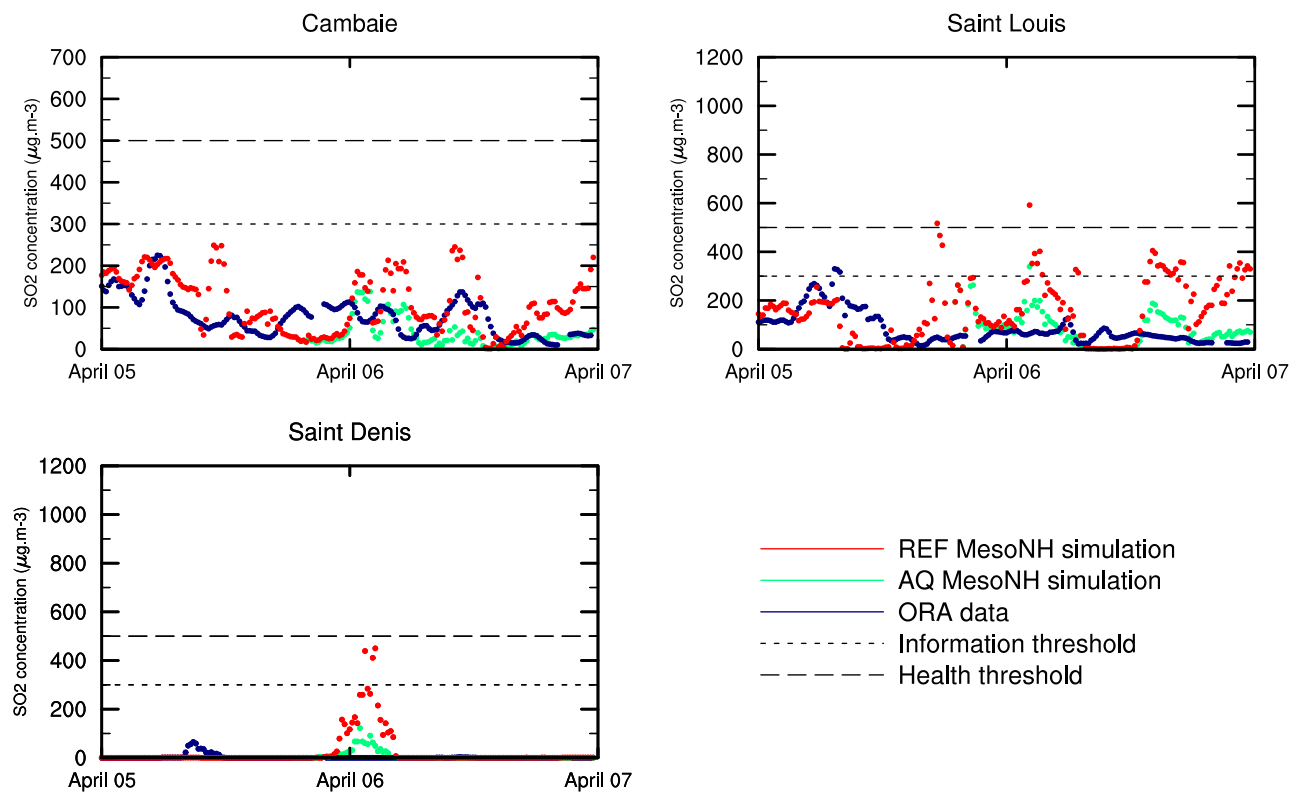
**Figure 9.** Cross section along  $\text{SO}_2$  plumes for April 3 (left) and April 6 (right) at 13 UTC. The cross sections are not in the same direction due to change of plume orientation. The left panel corresponds to blue line in Figure 1, the right panel to red line. Color filling corresponds to  $\text{SO}_2$  concentrations ( $\mu\text{g m}^{-3}$ ), isocontours values represent the upward velocity intensity ( $\text{m s}^{-1}$ ) generated by lava heat flow.



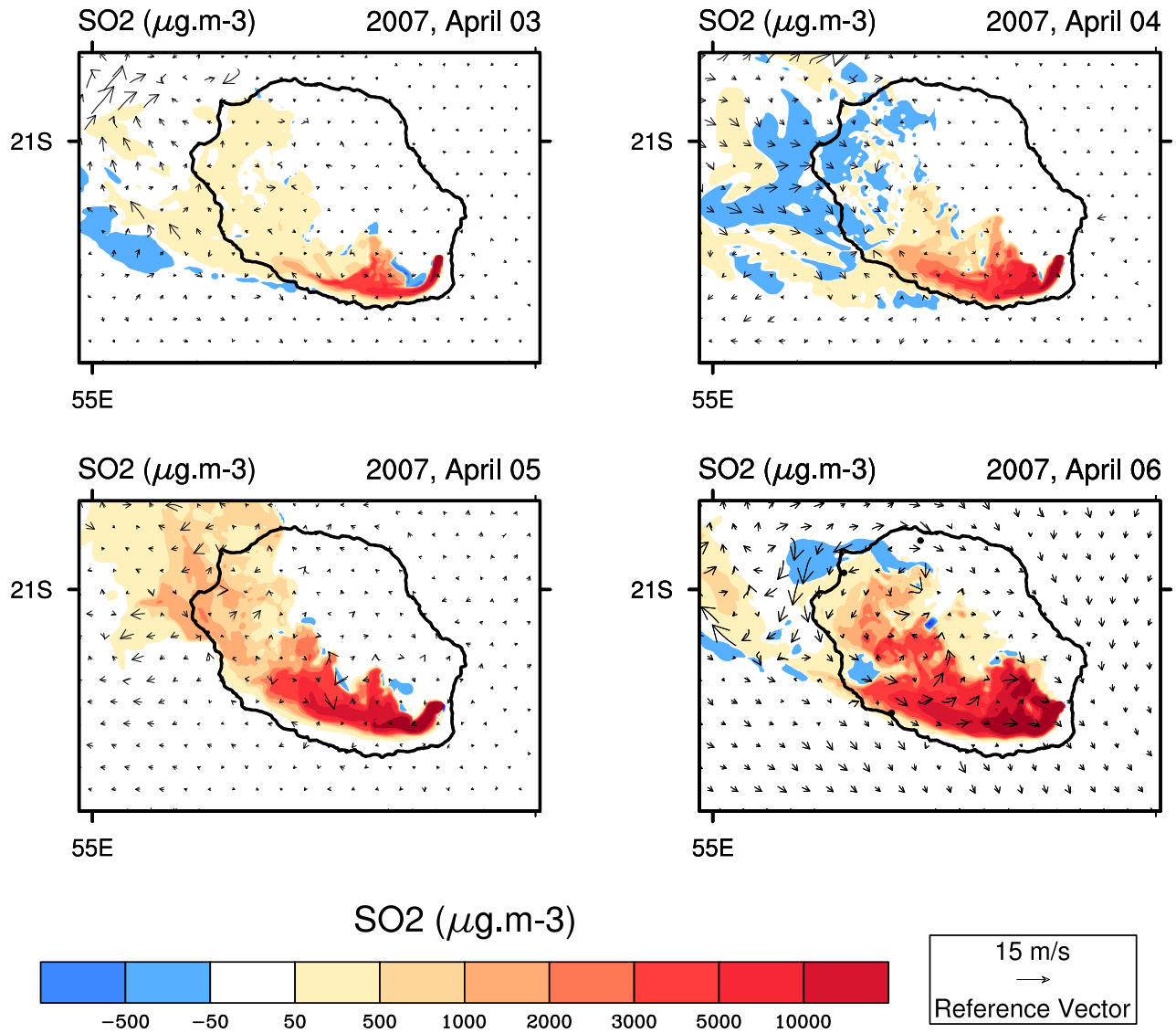
**Figure 10.** Accumulated rainfall given by MesoNH model between 2 and 7 April 2007. The numbers correspond to Meteo-France observations.



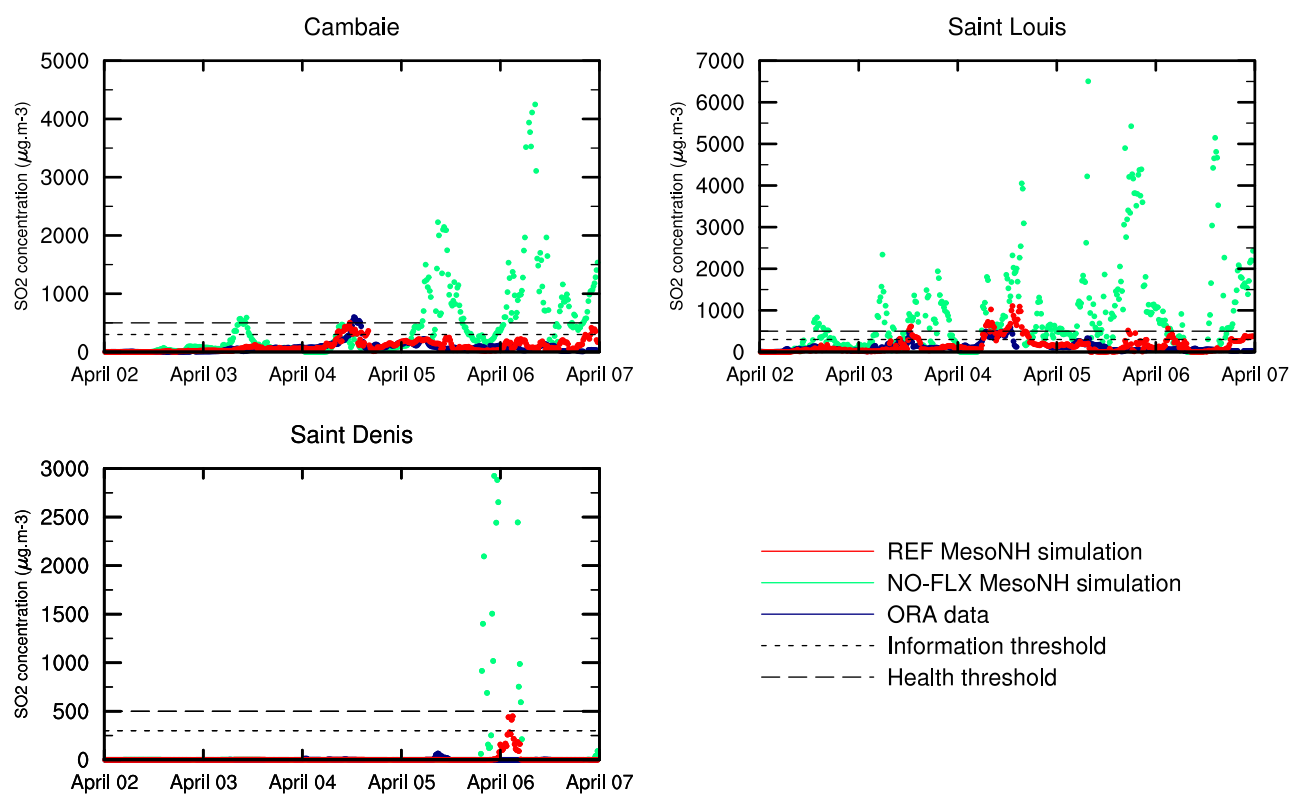
**Figure 11.** Difference of  $\text{SO}_2$  concentration between AQ and REF simulation the 6 April at 13 UTC at the surface (left panel) and in the plume (right panel).



**Figure 12.** Comparison surface  $\text{SO}_2$  concentration between between REF simulation and AQ simulation with observation providing by ORA.



**Figure 13.** SO<sub>2</sub> concentration difference at the surface between NO-FLX and REF simulations, for April 3,4,5 and 6 2007 at 13 UTC. The arrows represent the difference of the wind field between NO-FLX simulation and REF simulation.



**Figure 14.** Comparison of surface SO<sub>2</sub> concentration between NO-FLX simulation (green), REF simulation (red) and ORA measurements (blue). Thin dashed line is the public information threshold. Large dashed line is the health threshold.